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APPLICATION NO.	FILING DATE	FIRST NAMED INVENTOR	ATTORNEY DOCKET NO.	CONFIRMATION NO.
09/599,877	06/23/2000	Johan Lennerstrand	07691.0004	1424
7590 12/02/2003			EXAMINER	
PHILIP S. JOHNSON ONE JOHNSON & JOHNSON PLAZA NEW BRUNSWICK, NJ 08933-7003			PARKIN, JEFFREY S	
			ART UNIT	PAPER NUMBER
			1648	22
			DATE MAILED: 12/02/2003	

Please find below and/or attached an Office communication concerning this application or proceeding.

	Applicati n N .	Applicant(s)			
	09/599,877	LENNERSTRAND ET AL.			
Office Action Summary	Examin r	Art Unit			
	Jeffrey S. Parkin, Ph.D.	1648			
The MAILING DATE of this communication a Peri d for Reply	ppears on the c ver sheet	with the correspondence address			
A SHORTENED STATUTORY PERIOD FOR REF THE MAILING DATE OF THIS COMMUNICATION - Extensions of time may be available under the provisions of 37 CFR after SIX (6) MONTHS from the mailing date of this communication. - If the period for reply specified above is less than thirty (30) days, a re - If NO period for reply is specified above, the maximum statutory period. - Failure to reply within the set or extended period for reply will, by stat - Any reply received by the Office later than three months after the mail earned patent term adjustment. See 37 CFR 1.704(b). Status	N. 1.136(a). In no event, however, may reply within the statutory minimum of the dwill apply and will expire StX (6) Motute, cause the application to become	a reply be timely filed hirty (30) days will be considered timely. ONTHS from the mailing date of this communication. ABANDONED (35 U.S.C. § 133).			
1) Responsive to communication(s) filed on 22	September 2003.				
2a)⊠ This action is FINAL . 2b)□ Th	is action is non-final.				
	Since this application is in condition for allowance except for formal matters, prosecution as to the merits is closed in accordance with the practice under Ex parte Quayle, 1935 C.D. 11, 453 O.G. 213.				
Disposition of Claims					
4)⊠ Claim(s) <u>1-14,20 and 21</u> is/are pending in th	e application.				
4a) Of the above claim(s) is/are withdo					
5) Claim(s) is/are allowed.					
6)⊠ Claim(s) <u>1-14,20 and 21</u> is/are rejected.					
7) Claim(s) is/are objected to.					
8) Claim(s) are subject to restriction and	I/or election requirement.				
Application Papers					
9) The specification is objected to by the Exami	ner.				
10) The drawing(s) filed on is/are: a) a	ccepted or b) objected t	o by the Examiner.			
Applicant may not request that any objection to the	ne drawing(s) be held in abey	ance. See 37 CFR 1.85(a).			
Replacement drawing sheet(s) including the corre	ection is required if the drawir	ng(s) is objected to. See 37 CFR 1.121(d).			
11) The oath or declaration is objected to by the	Examiner. Note the attach	ed Office Action or form PTO-152.			
Priority under 35 U.S.C. §§ 119 and 120					
12) ☐ Acknowledgment is made of a claim for fore a) ☐ All b) ☐ Some * c) ☐ None of:		;. § 119(a)-(d) or (f).			
Certified copies of the priority docume Certified copies of the priority docume Copies of the certified copies of the priority docume Copies of the certified copies of the priority document in the International Bure * See the attached detailed Office action for a light for domain to the priority document in made of a plain for domain to the priority document in the priority d	ents have been received in riority documents have bee eau (PCT Rule 17.2(a)). ist of the certified copies no	en received in this National Stage ot received.			
13) Acknowledgment is made of a claim for dome since a specific reference was included in the 37 CFR 1.78.	first sentence of the specif	fication or in an Application Data Sheet.			
 a)	stic priority under 35 U.S.C	C. §§ 120 and/or 121 since a specific			
Attachment(s)					
1) Notice of References Cited (PTO-892) 2) Notice of Draftsperson's Patent Drawing Review (PTO-948)	5) 🔲 Notice o	w Summary (PTO-413) Paper No(s) of Informal Patent Application (PTO-152)			
3) Information Disclosure Statement(s) (PTO-1449) Paper No(s) 6) 🔲 Other:	-			

Serial No.: 09/599,877 Docket No.: 07691.0004

Applicants: Lennerstrand, J. and B. Larder Filing Date: 06/23/00

Response to Amendment

Status of the Claims

1. Acknowledgement is hereby made of receipt and entry of the amendment submitted 22 September, 2003, wherein claims 15-19 were canceled without prejudice or disclaimer and claims 1, 14, 20, and 21 amended. Claims 1-14, 20, and 21 are currently under examination.

35 U.S.C. § 112, Second Paragraph

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- 2. Claims 1-14, 20, and 21 stand rejected under 35 U.S.C. § 112, second paragraph, as being indefinite for failing to particularly point out and distinctly claim the subject matter which applicant regards as the invention.
- 3. The claims (1, 20, and 21) have been amended to include the phrase "the reaction products of substances" which remains vague and indefinite. The skilled artisan cannot readily ascertain the meaning of this phrase since the assay components are required for completion of the RT assay. Appropriate correction is required (i.e., providing a reaction well with the following assay components: I) at least one template for an HIV RT enzyme ...; providing a reaction well with the following assay reagents: I) at least one template for an HIV RT enzyme ...). Applicants are again directed toward pages 23 and 24 of the disclosure for suggestions in drafting appropriate claim language.
- 4. Claim 13 stands rejected under 35 U.S.C. § 112, second paragraph, as being indefinite for failing to particularly point out and distinctly claim the subject matter which applicant regards as the invention. The claim references various RT codons without providing a reference HIV-1 or HIV-2 isolate. Does the claim

reference the same codon in both viruses? Is the same mutation present in both HIV-1 and -2 drug-resistant RTs? Alternatively, does the claim simply refer to an HIV-1 RT mutation? correction is required.

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5. Claim 14 stands rejected under 35 U.S.C. § 112, second paragraph, as being indefinite for failing to particularly point out and distinctly claim the subject matter which applicant regards The claims have been amended to as the invention. reference to an "insertional mutation at nucleotide triplet encoding codon 69" which is still vague and indefinite since the precise nature and location of the mutation is not clearly set Perusal of the disclosure indicates the drug-resistant forth. forms of RT contain a single or multiple amino acid insertion between codons 69 and 70. Appropriate amendment of the claim language is required (i.e., wherein the HIV-1 mutant RT enzyme contains an amino acid insertion between codons 69 and 70).

35 U.S.C. § 112, First Paragraph

6. The following is a quotation of the first paragraph of 35 U.S.C. § 112:

The specification shall contain a written description of the invention, and of the manner and process of making and using it, in such full, clear, concise, and exact terms as to enable any person skilled in the art to which it pertains, or with which it is most nearly connected, to make and use the same and shall set forth the best mode contemplated by the inventor of carrying out his invention.

7. The previous rejection of claims 1-14, 20, and 21 under 35 30 U.S.C. § 112, first paragraph, as containing subject matter which was not described in the specification in such a way as to reasonably convey to one skilled in the relevant art that the inventor(s), at the time the application was filed, had possession 35

of the claimed invention, is hereby withdrawn in response to

applicants' arguments.

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35 U.S.C. § 103(a)

- 8. The following is a quotation of 35 U.S.C. § 103(a) which forms the basis for all obviousness rejections set forth in this Office action:
 - (a) A patent may not be obtained though the invention is not identically disclosed or described as set forth in section 102 of this title, if the differences between the subject matter sought to be patented and the prior art are such that the subject matter as a whole would have been obvious at the time the invention was made to a person having ordinary skill in the art to which said subject matter pertains. Patentability shall not be negatived by the manner in which the invention was made.

Subject matter developed by another person, which qualifies as prior art only under subsection (f) or (g) of section 102 of this title, shall not preclude patentability under this section where the subject matter and the claimed invention were, at the time the invention was made, owned by the same person or subject to an obligation of assignment to the same person.

- 9. This application currently names joint inventors. In considering patentability of the claims under 35 U.S.C. § 103(a), the examiner presumes that the subject matter of the various claims was commonly owned at the time any inventions covered therein were made absent any evidence to the contrary. Applicant is advised of the obligation under 37 C.F.R. § 1.56 to point out the inventor and invention dates of each claim that was not commonly owned at the time a later invention was made in order for the examiner to consider the applicability of 35 U.S.C. § 103© and potential 35 U.S.C. § 102(f) or (g) prior art under 35 U.S.C. § 103(a).
- 10. The factual inquiries set forth in *Graham et al. v. John Deere Company of Kansas City et al.; Calmar, Inc. v. Cook Chemical Company; Colgate-Palmolive Company v. Same,* 148 U.S.P.Q. 459 (U.S. Sup. Ct. 1966), that are applied for establishing a background for determining obviousness under 35 U.S.C. 103 are summarized as follows: 1) Determining the scope and contents of the prior art. 2) Ascertaining

the differences between the prior art and the claims at issue. 3) Resolving the level of ordinary skill in the pertinent art. 4) Considering objective evidence present in the application indicating obviousness or unobviousness.

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11. Claims 1-3, 5-12, 20, and 21 stand rejected under 35 U.S.C. § 103(a) as being unpatentable over Meyer et al. (1999) in view of Ekstrand et al. (1996). The claims are directed toward an HIV RT assay to assess the resistance of any given RT sample to treatment with an HIV RT inhibitor. The claims require a reaction well with the following components: (I) at least one template for an HIV RT enzyme; (ii) at least one primer; (iii) at least one detectable dNTP substrate; (iv) at least one HIV RT inhibitor; and (v) at least one ribonucleotide chosen from ATP and GTP, or at least one pyrophosphate. Additional steps recite comparative steps involving both the wildtype and mutant RTs.

As previously set forth, Meyer et al. (1999) provide an HIV RT enzymatic assay to examine mutant activity that employs at least one template, at least one primer, at least one RT inhibitor, and either ATP/GTP or pyrophosphate (see Experimental Procedures, p. The authors reported (p. 35, rt. col.) that "we describe an in vitro assay that reproduces the essential in vivo properties of HIV-1 RT containing the D67N, K70R, the AZT resistance mutants. and K219Q amino acid substitutions (designated 67/70/215/219 RT in this report) was much more efficient than WT RT at extending the primer past several potential termination sites in the presence of AZTTP when ATP was added to the reaction. of the AZTMP residue from the primer terminus to ATP to form dinucleoside polyphosphate and unblocked primer was enhanced in the 67/70/215/219 RT."

The authors also noted (see p. 35, last paragraph, rt. col.) that the "Addition of a ribonucleoside triphosphate (ATP) to the

reaction mixture provided an acceptor for the nucleotide-dependent primer unblocking activity in which the AZTMP residue from the chain-terminated primer was transferred to ATP to form Ap₄AZT, and the primer was shortened by one residue and was no longer blocked to elongation". The authors finally conclude (see p. 36, rt. col.) that "by adding ATP at concentrations likely to be present in intact cells, we have established an in vitro system that reflects the in vivo properties of the 67/70/215/219 mutant virus." The only limitation of this teaching is that it does not disclose an RT assay that employs a detectable dNTP.

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However, as previously set forth, Ekstrand et al. (1996) provide a non-radioactive reverse transcriptase assay that employs 5-bromodeoxyuridine 5'-triphosphate (BrdUTP) as the detectable dNTP (see Materials and Methods, p. 97). The assay described employs an alkaline phosphatase-conjugated anti-BrdU antibody and provides quantitative results. The authors note (see p. 104, last paragraph) "the present paper describes a simple, sensitive and non-radioactive RT assay with kinetic features similar to those observed when the natural dTTP substrate is used."

Therefore, it would have been prima facie obvious to one having ordinary skill in the art at the time the invention was made to utilize the detection format described by Ekstrand et al. (1996), in the RT assay provided by Meyer et al. (1999), since this provides a rapid, quantitative, and non-radioactive means for detecting the products of reverse transcription.

12. Applicants provide a declaration by Dr. Jochmans pursuant to 37 C.F.R. § 1.132 asserting that the claimed invention is unobvious in view of the prior art. The crux of the invention appears to be related to the use of a ribonucleotide (i.e., one of ATP or GTP) in the RT reaction mixture to facilitate the detection of drugresistant variant RTs. The inclusion of a ribonucleotide

apparently results in a more sensitive assay because it removes the block in polymerization stemming from the RT inhibitor. This is precisely the same format employed in the assay described by Meyer et al. (1999). In fact, Meyer and colleagues clearly stated that the inclusion of a ribonucleotide relieved the block in polymerization. Thus, contrary to the assertions of the declarant, the prior art appears to provide the crux of the claimed invention.

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It was further argued in the declaration that the prior art fails to teach the detection of multiple chain termination events This is precisely what Meyer and colleagues in a single well. The inclusion of the ribonucleotide relieves the block in polymerization thereby enabling one of ordinary skill in the art to detect multiple chain termination events by the mutant RT in the reaction well. The reaction conditions described by Meyer et al. (1999) are nearly identical to those described and claimed by applicants. The only deficiency in this teaching is its failure to describe the utilization of a labeled dNTP, such as BrdUTP. However, Ekstrand et al. (1996) provide a suitable label. Moreover, there was sufficient motivation to utilize this label in the assay of Meyer et al. (1999) and a reasonable expectation that the modified assay would be successful. Thus, the declarant's argument concerning this point is not persuasive.

Applicants have previously argued that sufficient motivation and a reasonable expectation of success were not present in the prior art. These arguments were clearly not persuasive in view of the prior art. Moreover, as previously set forth, the Examiner recognizes that references cannot be arbitrarily combined and that there must be some reason why one skilled in the art would be motivated to make the proposed combination of primary and secondary references. In re Nomiya, 184 U.S.P.Q. 607 (C.C.P.A. 1975). However, there is no requirement that a motivation to make the modification be expressly articulated. The test for combining

references is what the combination of disclosures taken as a whole would suggest to one of ordinary skill in the art. In re McLaughlin, 170 U.S.P.Q. 209 (C.C.P.A. 1971). References are evaluated by what they suggest to one versed in the art, rather than by their specific disclosures. In re Bozek, 163 U.S.P.Q. 545 (C.C.P.A. 1969). As set forth supra, both the motivation and a reasonable expectation of success were present in the prior art. One of ordinary skill in the art would have had sufficient motivation to utilize the detection format described by Ekstrand et al. (1996), in the RT assay provided by Meyer et al. (1999), since this would provide a rapid, quantitative, and non-radioactive means for detecting the products of reverse transcription.

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- Claim 4 stands rejected under 35 U.S.C. § 103(a) as being 13. unpatentable over Meyer et al. (1999) in view of Ueno et al. The content of Meyer et al. (1999) is disclosed in the Meyer and colleagues do not describe the preceding paragraph. utilization of an art-recognized RT activity label such as a radioactive dNTP, although a labeled primer was employed. However, Ueno et al. (1995) describe standard HIV RT assays that employ artrecognized labels such as radioactive labeled dNTPs (see pp. 23605-23606, EXPERIMENTAL PROCEDURES, Materials and Product Analysis). Therefore, it would have been prima facie obvious to one having ordinary skill in the art at the time the invention was made to utilize a radiolabeled dNTP, as taught by Ueno et al. (1995), in the assay of Meyer et al. (1999), since this represents a standard and art-recognized means for detecting RT reaction products. Applicants' arguments are not convincing as noted in the preceding paragraph.
- 14. Claims 13 and 14 stand rejected under 35 U.S.C. § 103(a) as being unpatentable over Meyer et al. (1999) in view of Ekstrand et

al. (1996), as applied supra to claims 1-3, 5-12, 20, and 21, and further in view of Larder et al. (1999a, 1999b). The combination of references employed supra do not disclose the use of HIV RT mutants carrying mutations at amino acid positions 67, 69, and 70, or an insertion between amino acids 69 and 70. However, both Larder et al. (1999a, 1999b) publications disclose that HIV-1 RT resistant variants, particularly MNR variants, carry mutations at amino acids 67, 69, and 70, and between amino acids 69 and 70. Therefore, it would have been prima facie obvious to one having ordinary skill in the art at the time the invention was made to utilize the HIV-1 RT mutants described by Larder et al. (1999a, 1999b), in the reverse transcriptase assay suggested by Ekstrand et al. (1996) and Meyer et al. (1999). One of ordinary skill in the art would have been motivated to include these mutant forms of the RT since they naturally develop during the course of antiviral therapy. Applicants' arguments are not convincing for the reasons of record set forth supra.

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Claims 1-3, 5-12, 20, and 21 stand rejected under 35 U.S.C. 15. § 103(a) as being unpatentable over Arion et al. (1998) in view of Ekstrand et al. (1996). Arion et al. (1999) provides an HIV RT enzymatic assay to examine mutant activity that employs a template, primer, RT inhibitor, and pyrophosphate (see p. 15910, MATERIALS AND METHODS, Analysis of Chain Termination of RT-Catalyzed DNA The authors suggested (see p. 15908, ABSTRACT) that Synthesis). "HIV-1 resistance to AZT results from the selectively decreased binding of AZTTP and the increased pyrophosphorolytic cleavage of chain-terminated viral DNA by the mutant RT at physiological pyrophosphate levels, resulting in a net decrease in chain termination. The increased processivity of viral DNA synthesis may be important to enable facile HIV replication in the presence of AZT, by compensating for the increased reverse reaction rate."

This teaching does not disclose an RT assay that employs a detectable dNTP.

However, Ekstrand et al. (1996) provide a non-radioactive reverse transcriptase assay that employs 5-bromodeoxyuridine 5'-triphosphate (BrdUTP) as the detectable dNTP (see Materials and Methods, p. 97). The assay described employs an alkaline phosphatase-conjugated anti-BrdU antibody and provides quantitative results. The authors note (see p. 104, last paragraph) "the present paper describes a simple, sensitive and non-radioactive RT assay with kinetic features similar to those observed when the natural dTTP substrate is used."

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Therefore, it would have been prima facie obvious to one having ordinary skill in the art at the time the invention was made to utilize the detection format described by Ekstrand et al. (1996), in the RT assay provided by Arion et al. (1998), since this provides a rapid, quantitative, and non-radioactive means for detecting the products of reverse transcription.

16. Applicants' arguments set forth in the declaration of Dr. Jochmans were addressed above. Applicants previously argued that both sufficient motivation and a reasonable expectation of success were not present in the prior art is not convincing. arguments are clearly not persuasive in view of the prior art and Moreover, the Examiner knowledge of the skilled artisan. recognizes that references cannot be arbitrarily combined and that there must be some reason why one skilled in the art would be motivated to make the proposed combination of primary and secondary In re Nomiya, 184 U.S.P.Q. 607 (C.C.P.A. 1975). references. However, there is no requirement that a motivation to make the modification be expressly articulated. The test for combining references is what the combination of disclosures taken as a whole would suggest to one of ordinary skill in the art.

McLaughlin, 170 U.S.P.Q. 209 (C.C.P.A. 1971). References are evaluated by what they suggest to one versed in the art, rather than by their specific disclosures. In re Bozek, 163 U.S.P.Q. 545 (C.C.P.A. 1969). As set forth supra, both the motivation and a reasonable expectation of success were present in the prior art. One of ordinary skill in the art would have had sufficient motivation to utilize the detection format described by Ekstrand et al. (1996), in the RT assay provided by Arion et al. (1998), since this provides a rapid, quantitative, and non-radioactive means for detecting the products of reverse transcription.

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- Claim 4 stands rejected under 35 U.S.C. § 103(a) as being 17. unpatentable over Arion et al. (1998) in view of Ueno et al. The content of Arion et al. (1998) is disclosed in the preceding paragraph. Meyer and colleagues do not describe the utilization of an art-recognized RT activity label such as a radioactive dNTP, although a labeled primer was employed. However, Ueno et al. (1995) describe standard HIV RT assays that employ artrecognized labels such as radioactive labeled dNTPs (see pp. 23605-23606, EXPERIMENTAL PROCEDURES, Materials and Product Analysis). Therefore, it would have been prima facie obvious to one having ordinary skill in the art at the time the invention was made to utilize a radiolabeled dNTP, as taught by Ueno et al. (1995), in the assay of Arion et al. (1998), since this represents a standard and art-recognized means for detecting RT reaction products. Applicants' arguments are not convincing for the reasons set forth in the preceding paragraph.
- 18. Claims 13 and 14 stand rejected under 35 U.S.C. § 103(a) as being unpatentable over Arion et al. (1998) in view of Ekstrand et al. (1996), as applied supra to claims 1-3, 5-12, 20, and 21, and further in view of Larder et al. (1999a, 1999b). The combination

of references employed supra do not disclose the use of HIV RT mutants carrying mutations at amino acid positions 67, 69, and 70, or an insertion between amino acids 69 and 70. However, both Larder et al. (1999a, 1999b) publications disclose that HIV-1 RT resistant variants, particularly MNR variants, carry mutations at amino acids 67, 69, and 70, and between amino acids 69 and 70. Therefore, it would have been prima facie obvious to one having ordinary skill in the art at the time the invention was made to utilize the HIV-1 RT mutants described by Larder et al. (1999a, 1999b), in the reverse transcriptase assay suggested by Ekstrand et al. (1996) and Arion et al. (1998). One of ordinary skill in the art would have been motivated to include these mutant forms of the RT since they naturally develop during the course of antiviral therapy. Applicants' arguments are not convincing for the reasons of record set forth supra.

Finality of Office Action

19. THIS ACTION IS MADE FINAL. Applicant is reminded of the extension of time policy as set forth in 37 C.F.R. § 1.136(a). A SHORTENED STATUTORY PERIOD FOR RESPONSE TO THIS FINAL ACTION IS SET TO EXPIRE THREE MONTHS FROM THE DATE OF THIS ACTION. IN THE EVENT A FIRST RESPONSE IS FILED WITHIN TWO MONTHS OF THE MAILING DATE OF THIS FINAL ACTION AND THE ADVISORY ACTION IS NOT MAILED UNTIL AFTER THE END OF THE THREE-MONTH SHORTENED STATUTORY PERIOD, THEN THE SHORTENED STATUTORY PERIOD WILL EXPIRE ON THE DATE THE ADVISORY ACTION IS MAILED, AND ANY EXTENSION FEE PURSUANT TO 37 C.F.R. § 1.136(a) WILL BE CALCULATED FROM THE MAILING DATE OF THE ADVISORY ACTION. IN NO EVENT WILL THE STATUTORY PERIOD FOR RESPONSE EXPIRE LATER THAN SIX MONTHS FROM THE DATE OF THIS FINAL ACTION.

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UPERVISORY PATENT EXAMINER
TECHNOLOGY CENTER 1600

Correspondence

Correspondence related to this application may be submitted to Group 1600 by facsimile transmission. The faxing of such papers must conform with the notice published in the Official Gazette, 1096 OG 30 (November 15, 1989). Official communications should be directed toward the following Group 1600 fax number: (703) 872-9306. Any inquiry concerning this communication should be directed to Jeffrey S. Parkin, Ph.D., whose telephone number is (703) 308-2227. The examiner can normally be reached Monday through Thursday from 8:30 AM to 6:00 PM. A message may be left on the examiner's If attempts to reach the examiner are voice mail service. unsuccessful, the examiner's supervisors, Laurie Scheiner or James Housel, can be reached at (703) 308-1122 or (703) 308-4027, respectively. Any inquiry of a general nature or relating to the status of this application should be directed to the Group 1600 receptionist whose telephone number is (703) 308-0196.

Respectfully,

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Jeffrey S. Parkin, Ph.D.

Patent Examiner
Art Unit 1648

30 November, 2003